Advanced Regression: Random effects and hierarchical models

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4. Random effects

1. Random effect model with random intercept

$$y_i = (\alpha_0 + u_k) + \beta x_i + \epsilon_i,$$

where $u_k \sim N(0, \sigma_u^2)$.

2. Random effects model on both, the intercept and the

$$y_i = (\alpha_0 + u_k) + (\beta + w_k) x_i + \epsilon_i,$$
 where $w_k \sim N(0, \sigma_w^2).$

Random intercept

1. Random effect model with random intercept

 $y_i = (\alpha_0 + u_k) + \beta x_i + \epsilon_i, = \alpha_0 + \beta x_i + u_k + \epsilon_i,$ where α_0 is the intercept and β the regression coefficient.

- There are two distinct error terms
 - 1. Group-specific error $u_k \sim N(0, \sigma_u^2)$ 2. Individual-specific error $\epsilon_i \sim N(0, \sigma^2)$
- Note that u_k and ϵ_i are independent of each other.

Group effects are also called random effects:

- 1. Random effect for the intercept $u_k \sim N(0, \sigma_u^2)$
- 2. Random effect for the slope $w_k \sim N(0, \sigma_w^2)$

Random effect model with random intercept

Interpretation of random intercept α_k :

$$\alpha_k = (\alpha_0 + u_k)$$

- α_0 is the global intercept
- u_k group-level variations around the global intercept

This is equivalent to assuming α_k is a random variable that follows a Normal distribution

$$\alpha_k \sim N(\alpha_0, \sigma_u^2)$$

Random effect model with random intercept

Multi-level interpretation (two levels of variability):

1. **First level.** Defined on the individual level for observation i = 1, ..., n, similar to a standard linear regression

$$y_i = \alpha_k + \beta x_i + \epsilon_i$$

2. **Second level**. But the intercept is not fixed, it is a random variable

$$\alpha_k \sim N(\alpha_0, \sigma_u^2)$$

Random effect model with random intercept

Assumptions:

- Slope of regression line is the same across all groups. Each group has a different intercept (α_k) .
- But $\alpha_k \sim N(\alpha_0, \sigma_u^2)$ has now a common distribution which is estimated from **all observations**, and not just from the observations in a specific group as in fixed effects.
- We pool information across groups.

Consequences:

- We control for group characteristics by including the group-specific intercept.
- Number of group-specific parameters to estimate is much smaller than in the fixed effect models (σ_u^2 vs k intercepts).

(Restricted) Maximum Likelihood estimation of random effect

$$y_i = \alpha_0 + \beta x_i + u_k + \epsilon_i$$

Parameters to estimate are

- α_0, β intercept and regression coefficient σ_u^2, σ^2 variance components

Maximum Likelihood estimation is based on the Normal distribution of u_k and ϵ_i

- ML estimate for σ_u^2 requires subtracting 2 empirical estimates of variance \to ML estimates for σ_u^2 can be negative.
- Restricted Maximum Likelihood (REML): Imposes positivity constraints on the variance estimates.

Random effects in R: nlme::lme()

Implementations of Restricted Maximum Likelihood (REML) in R:

- lmer function in the lme4 package
- lme function in the nlme package

Focus here is on nlme::lme(fixed, random, data):

- fixed. Formula $y \sim x$
- random. Formula ~ 1 | factor
- data. Dataset to use

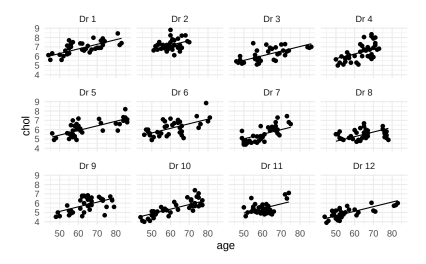
R: Random intercept using lme()

```
model_random_intercept <- lme(chol ~ age, random = ~ 1 | doctor, data = data_chol)</pre>
summary(model_random_intercept)
Linear mixed-effects model fit by REML
 Data: data_chol
      AIC
             BIC
                     logLik
  828.697 845.035 -410.3485
Random effects:
Formula: ~1 | doctor
        (Intercept) Residual
StdDev:
          0.6347908 0.5764246
Fixed effects: chol ~ age
                Value Std.Error DF t-value p-value
(Intercept) 2.9060357 0.26477408 428 10.97553
           0.0495831 0.00306279 428 16.18888
                                                    0
Correlation:
    (Intr)
age -0.714
Standardized Within-Group Residuals:
                   Q1
                             Med
                                         Q3
                                                   Max
-2.7850636 -0.7012544 -0.1419474 0.6536599 3.0850909
Number of Observations: 441
Number of Groups: 12
```

R: Random intercept using lme()

```
data_chol |>
  mutate(
    doctor_name = factor(str_c("Dr ", doctor), levels = str_c("Dr ", 1:12)),
    .fitted = fitted(model_random_intercept)
) |>
```

```
ggplot(aes(x = age)) +
geom_point(aes(y = chol)) +
geom_line(aes(y = .fitted)) +
facet_wrap(~doctor_name) +
theme_minimal()
```



Random effect model and variance partition

Variance decomposition for observation i in group k

$$var(y_i) = var(u_k + \epsilon_i) \tag{1}$$

$$= \mathrm{var}(u_k) + \mathrm{var}(\epsilon_i) + 2\mathrm{cov}(u_k, \epsilon_i) \tag{2}$$

$$= \sigma_u^2 + \sigma^2 + 0 \tag{3}$$

Further we can look at the covariance of observations

• i and i' within group k

$$\mathrm{cov}(y_i,y_{i'}) = \mathrm{cov}(u_k + \epsilon_i, u_k + \epsilon_{i'}) = \sigma_u^2$$

• i and i' from different groups k and k'

$$cov(y_i, y_{i'}) = cov(u_k + \epsilon_i, u_{k'} + \epsilon_{i'}) = 0$$

Random effect model and variance partition

Variability between and within groups. Intra-class correlation coefficient ρ

$$\rho = \operatorname{cor}(y_i, y_{i'}) = \frac{\operatorname{cov}(y_i, y_{i'})}{\sqrt{\operatorname{var}(y_i)\operatorname{var}(y_{i'})}} = \frac{\sigma_u^2}{\sigma_u^2 + \sigma^2}$$

Interpretation:

- Intra-class correlation coefficient ρ is the correlation between two observations i and i' in the same group.
- It is the ratio of between-group variance σ_u^2 over the total variance
- If $\rho \to 0$ there is little variation explained by the grouping and we might consider a model without the random effect.
- Any restrictions?

Variance partition in R

VarCorr(model_random_intercept)

$$\rho = \frac{\sigma_u^2}{\sigma_u^2 + \sigma^2} = \frac{0.6347908^2}{0.6347908^2 + 0.5764246^2} \approx 0.54$$

Interpretation:

- There is substantial evidence for between-group heterogeneity.
- More than half of the total variance can be explained by the between-group variance.
- It is beneficial to include the random effects on the intercept.

Random effect model with random intercept and random slope

Random effects model on both the intercept and the slope

$$y_i = (\alpha_0 + \mathbf{u_k}) + (\beta + \mathbf{w_k})x_i + \epsilon_i$$

- There are three distinct error terms
 - 1. Group-specific error of the intercept

$$u_k \sim N(0, \sigma_u^2)$$

2. Group-specific error of the regression slope

$$w_k \sim N(0, \sigma_w^2)$$

3. Individual-specific error

$$\epsilon_i \sim N(0, \sigma^2)$$

• Note that u_k and w_k are correlated and independent of ϵ_i .

Random effect model with random intercept and random slope

Assumptions:

- Each group has a different intercept $(\alpha_k = \alpha_0 + u_k)$ and a different regression slope $(\beta_k = \beta + w_k)$.
- We allow for correlation between α_k and β_k .
- Both, $\alpha_k \sim N(\alpha_0, \sigma_u^2)$ and $\beta_k \sim N(\beta, \sigma_w^2)$ have a common distribution which is estimated from **all observations**, and not just from the observations in a given group as in fixed effects.
- We pool information across groups.

Consequences:

- Including a random slope can be interpreted as creating an interaction between the group and the strength of association.
- We only have three additional parameters in the model: σ_u^2, σ_w^2 and $cor(\sigma_u, \sigma_w)$.

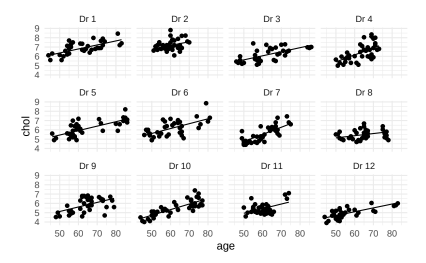
R: Random intercept and slope using lme()

```
model_random_slope <- lme(chol ~ age, random = ~ 1 + age | doctor, data = data_chol)</pre>
summary(model_random_slope)
Linear mixed-effects model fit by REML
 Data: data_chol
       AIC
               BIC
                       logLik
  821.9886 846.4956 -404.9943
Random effects:
Formula: ~1 + age | doctor
Structure: General positive-definite, Log-Cholesky parametrization
           StdDev
(Intercept) 1.28163949 (Intr)
           0.01771589 -0.872
Residual
           0.55997507
Fixed effects: chol ~ age
                Value Std.Error DF t-value p-value
(Intercept) 2.8791744 0.4215204 428 6.830451
           0.0500704 0.0060597 428 8.262823
Correlation:
    (Intr)
age -0.901
Standardized Within-Group Residuals:
       Min
                             Med
                   Q1
                                         QЗ
                                                   Max
-2.8523390 -0.6664557 -0.1141924 0.6206844 3.0809629
Number of Observations: 441
Number of Groups: 12
```

R: Random intercept and slope using lme()

```
data_chol |>
  mutate(
```

```
doctor_name = factor(str_c("Dr ", doctor), levels = str_c("Dr ", 1:12)),
    .fitted = fitted(model_random_slope)
) |>
    ggplot(aes(x = age)) +
    geom_point(aes(y = chol)) +
    geom_line(aes(y = .fitted)) +
    facet_wrap(~doctor_name) +
    theme_minimal()
```



Variables on individual and group level

When considering variables or predictors we need to distinguish:

- Individual-level variables
- **Group-level variables** that are the same for all observations in a group

GP example:

- Individual-level variables: Age and sex of patient
- Group-level variables: Age of doctor

```
# A tibble: 6 x 6
  chol doctor
                      bmi agedoc
                age
  <dbl> <dbl> <dbl> <dbl> <
                          <dbl> <dbl>
1 7.13
                 54 27.4
            1
                              55
                                     0
2 7.7
            1
                 55 29.1
                              55
                                     0
3 7.3
                 56 27.9
                              55
                                     0
            1
4 6.89
            1
                 71 26.7
                              55
                                     1
5 6.9
                 72 26.7
                              55
6 7.9
                 73 29.7
                              55
```

Variables on individual and group level

```
y_i = (\alpha_0 + u_k) + (\beta + w_k)x_i + \frac{\gamma x_q}{\epsilon_i} + \epsilon_i
```

```
model_random_cov <- lme(chol ~ age + agedoc, random = ~ 1 + age | doctor, data = data_chol)
summary(model_random_cov)</pre>
```

```
Linear mixed-effects model fit by REML
 Data: data_chol
       AIC
                BIC
                       logLik
  815.6956 844.2712 -400.8478
Random effects:
 Formula: ~1 + age | doctor
Structure: General positive-definite, Log-Cholesky parametrization
            StdDev
                       Corr
(Intercept) 1.10559524 (Intr)
            0.01775586 -0.951
age
Residual
           0.55992053
Fixed effects: chol ~ age + agedoc
                 Value Std.Error DF
                                       t-value p-value
(Intercept) -2.7897788 1.1824050 428 -2.359410 0.0188
             0.0501492 0.0060673 428 8.265423 0.0000
age
             0.1280030 0.0253576 10 5.047908 0.0005
agedoc
Correlation:
       (Intr) age
      -0.303
age
agedoc -0.948 -0.004
```

```
Standardized Within-Group Residuals:
```

```
Min Q1 Med Q3 Max -2.8426263 -0.6665146 -0.1046213 0.6206861 3.0950474
```

Number of Observations: 441

Number of Groups: 12

Model comparison

Likelihood-ratio test for nested models:

- Models must have the same fixed effects. Does not work with group-level covariates.
- Model with smaller log likelihood is better (better model fit).

Akaike information criterion (AIC):

• Model with the smaller AIC is better (less information loss).

Model comparison

GP example:

- Model A (Random intercept) model_random_intercept
 = lme(chol ~ age, random = ~ 1 | doctor, data =
 data_chol)
- Model B (Random intercept and slope) model_random_slope
 = lme(chol ~ age, random = ~ 1 + age | doctor,
 data = data_chol)
- Model C (Random intercept and slope and group covariate) model_random_cov = lme(chol ~ age + agedoc, random = ~ 1 + age | doctor, data = data_chol)

Model comparison

Likelihood-ratio test for nested models:

```
anova(model_random_intercept, model_random_slope)
```

```
Model df AIC BIC logLik Test L.Ratio model_random_intercept 1 4 828.6970 845.0350 -410.3485 model_random_slope 2 6 821.9886 846.4956 -404.9943 1 vs 2 10.7084 p-value model_random_intercept model_random_slope 0.0047
```

Akaike information criterion (AIC):

```
anova(model_random_slope, model_random_cov)
```

Warning in anova.lme(model_random_slope, model_random_cov): fitted objects with different fixed effects. REML comparisons are not meaningful.

```
Model df AIC BIC logLik Test L.Ratio p-value model_random_slope 1 6 821.9886 846.4956 -404.9943 model_random_cov 2 7 815.6956 844.2712 -400.8478 1 vs 2 8.292926 0.004
```

Generalised linear mixed models

- Generalised Linear Mixed models (GLMM) can be used to adapt linear mixed models to outcomes that do not follow a Normal distribution.
- The package lme4 includes the function glmer that can fit GLMMs.

```
glmer(formula = y \sim x + (1 + x | factor), family = gaussian)
```

Take away: Fixed and random effects

- Fixed effect models can account for group structure but many parameters need to be estimated and no information is shared between groups.
- Random effect models treat group-specific parameters as random variables.
- Instead of estimating one parameter for each group, random effect models only estimate the distribution parameter of the random variable.
- Thus, they pool information across groups.
- The intra-class coefficient gives a measure of how relevant the group structure is.
- Implementation in R: lme() function in the nlme package.
- Models including both, fixed and random effects, are often called linear mixed models.

Questions?